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FILE CONTENT: 1840 - 13 Oct 2007 VOL 147 ISS 17

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Some CASREACT records are derived from the ZIC/VINITI database (1974-1999) provided by InfoChem, INPI data prior to 1986, and Biotransformations database compiled under the direction of Professor Dr. Klaus Kieslich.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d que L1 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

L3 2 SEA FILE=CASREACT SSS FUL L1 (6 REACTIONS)

=> d 13 1-2 ibib abs fcrd

L3 ANSWER 1 OF 2 CASREACT COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 140:111409 CASREACT

TITLE: A novel process to prepare pioglitazone via several

novel intermediates.

INVENTOR(S): Pandey, Bipin; Lohray, Vidya Bhushan; Lohray, Braj

Bhushan

PATENT ASSIGNEE(S): Cadila Healthcare Limited, India

SOURCE: PCT Int. Appl., 74 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND						DATE			A.	PPLI	CATI	ON N	ο.	DATE					
									-										
WO	WO 2004007490 A2						20040122 WO 2003-IN241 20030715												
WO	WO 2004007490 A3					2004	0325												
	W:	AE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,		
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,		
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,		
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	ΝZ,	OM,		
		PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ТJ,	TM,	TN,		
		TР	ידיים	T_{2}	TΤΔ	TIC	TIC	117	VC	VIN	VII	7. D	7.M	7.W					

GI

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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
              KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
              FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
              BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                               IN 2002-MU648
     IN 2002MU00648
                              20040424
                                                                  20020716
                        Α
     AU 2003272072
                                               AU 2003-272072
                                                                  20030715
                         A1
                              20040202
     EP 1521753
                         A2
                              20050413
                                               EP 2003-753913
                                                                  20030715
     EP 1521753
                        B1
                              20070905
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
                              20070915
                                               AT 2003-753913
                                                                  20030715
     AT 372336
     US 2006167061
                              20060727
                                               US 2005-520166
                                                                  20051004
                         A1
PRIORITY APPLN. INFO.:
                                               IN 2002-MU648
                                                                  20020716
                                               WO 2003-IN241
                                                                  20030715
                          MARPAT 140:111409
OTHER SOURCE(S):
```

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- The present invention discloses a novel and general process to prepare AB various pyridine substituted 5-[4-[2-(alkyl substituted pyridyl)ethoxy]benzyl]-2,4-thiazolidinone derivs. of general formula I [R = alkyl], and their pharmaceutically acceptable salts. The present invention especially provides a novel process to prepare pioglitazone hydrochloride [R = 5-ethyl], via novel intermediates, i.e. II and III. This process involves lesser number of steps with high yields and uses key solid intermediates, which are operationally simple, and therefore offers opportunities for better com. viability.

RX(1) OF 92 - REACTION DIAGRAM NOT AVAILABLE

ANSWER 2 OF 2 CASREACT COPYRIGHT 2007 ACS on STN

126:42525 CASREACT ACCESSION NUMBER:

Synthesis and Biological Activity of Metabolites of TITLE:

the Antidiabetic, Antihyperglycemic Agent Pioglitazone

Tanis, Steven P.; Parker, Timothy T.; Colca, Jerry R.; AUTHOR (S):

Fisher, Roberta M.; Kletzein, Rolf F.

Department of Discovery Chemistry, Pharmacia and CORPORATE SOURCE:

Upjohn Inc., Kalamazoo, MI, 49001, USA

SOURCE:

Journal of Medicinal Chemistry (1996), 39(26),

5053-5063

CODEN: JMCMAR; ISSN: 0022-2623

American Chemical Society

DOCUMENT TYPE: Journal English LANGUAGE:

GI

PUBLISHER:

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- The authors have developed improved syntheses of pioglitazone metabolites AB I, II, III, and IV and the putative metabolite ketone V. These entities have been compared in the KKAy mouse model of human type-II diabetes to pioglitazone. Ketone V has proven to be the most potent of these thiazolidinediones in this in vivo assay. When I-V were compared in vitro in the 3T3-L1 cell line to pioglitazone for their ability to augment insulin-stimulated lipogenesis, V was again the most potent compound with I,

II, and IV roughly equivalent to pioglitazone. These data suggest that metabolites I, II, and IV are likely to contribute to the pharmacol. activity of pioglitazone, as had been previously reported for ciglitazone..

RX(4) OF 114 - REACTION DIAGRAM NOT AVAILABLE
REFERENCE COUNT: 37 THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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FILE COVERS 1907 - 17 Oct 2007 VOL 147 ISS 17 FILE LAST UPDATED: 16 Oct 2007 (20071016/ED)

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http://www.cas.org/infopolicy.html

Structure attributes must be viewed using STN Express query preparation. L5 STR

G1 OH, X, O

Structure attributes must be viewed using STN Express query preparation.

L6 109 SEA FILE=REGISTRY SSS FUL L4
L7 22 SEA FILE=REGISTRY SSS FUL L5
L8 26 SEA FILE=CAPLUS L6 AND L7

=> d 18 1-26 ibib abs hitstr

L8 ANSWER 1 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2007:1091018 CAPLUS

TITLE:

Combination therapy formulations comprising

thiazolidinedione analogues, glucocorticoid agonist, and nonsteroidal antiinflammatory drugs for treating

inflammatory disease

INVENTOR(S):

Colca, Gerald R.; Kletzien, Rolf F.

PATENT ASSIGNEE(S):

Metabolic Solutions Development Company, USA

SOURCE:

PCT Int. Appl., 48pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	KIND DATE					APPLICATION NO.						DATE					
					-												
WO 2007109088				A2		20070927		1	WO 2007-US6508					20070314			
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	KΡ,	KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	MG,	MK,	MN,	
	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RS,	
	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	sv,	SY,	ТJ,	TM,	TN,	TR,	TT,	TZ,	
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RW	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	
	IS,	IT,	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	
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	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	
	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM										

PRIORITY APPLN. INFO.:

US 2006-782972P P 20060316

The present invention relates to thiazolidinedione analogs that are useful for treating inflammatory disease. In general, the invention relates to pharmaceutical compns. comprising a combination of glucocorticoid agonists and insulin sensitizers that have reduced activation of the nuclear transcription factor PPARy. Thus, pharmaceutical composition including a thiazolidinedione analogs can be produced by tableting between 1 mg to 200 mg thiazolidinedione analog, CM-cellulose or carmellose, magnesium stearate, hydroxypropyl cellulose, and lactose monohydrate.

IT 146062-49-9 950696-94-3 950696-95-4 950696-96-5 950696-97-6 950696-98-7

CN

950696-99-8 950697-00-4 950697-01-5 950697-03-7 950697-05-9 950697-07-1

950697-09-3

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (combination therapy formulations comprising thiazolidinedione analogs, glucocorticoid agonist, and nonsteroidal antiinflammatory drugs for treating inflammatory disease)

146062-49-9 CAPLUS RN

2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2oxoethoxy]phenyl]methyl] - (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

Et

RN 950696-94-3 CAPLUS

2,4-Thiazolidinedione, 5-[[4-[(2R)-2-(5-ethyl-2-pyridinyl)-2-CN hydroxyethoxy]phenyl]methyl] - (CA INDEX NAME)

Absolute stereochemistry.

950696-95-4 CAPLUS RN

Ēt

ANSWER 2 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2007:1090689 CAPLUS

TITLE:

Thiazolidinedione analogues for treating hypertension

INVENTOR(S):

Colca, Gerard R.; Kletzien, Rolf F.

PATENT ASSIGNEE(S):

Metabolic Solutions Development Company, USA

PCT Int. Appl., 57pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 2007109037	A2 20070	927 WO 2007-US6385	20070314
W: AE, AG, AL,	AM, AT, AU,	AZ, BA, BB, BG, BR, BW,	BY, BZ, CA, CH,
CN, CO, CR,	CU, CZ, DE,	DK, DM, DZ, EC, EE, EG,	ES, FI, GB, GD,
GE, GH, GM,	GT, HN, HR,	HU, ID, IL, IN, IS, JP,	KE, KG, KM, KN,
KP, KR, KZ,	LA, LC, LK,	LR, LS, LT, LU, LY, MA,	MD, MG, MK, MN,
MW, MX, MY,	MZ, NA, NG,	NI, NO, NZ, OM, PG, PH,	PL, PT, RO, RS,
RU, SC, SD,	SE, SG, SK,	SL, SM, SV, SY, TJ, TM,	TN, TR, TT, TZ,
ÚA, UG, US,	UZ, VC, VN,	ZA, ZM, ZW	
RW: AT, BE, BG,	CH, CY, CZ,	DE, DK, EE, ES, FI, FR,	GB, GR, HU, IE,
IS, IT, LT,	LU, LV, MC,	MT, NL, PL, PT, RO, SE,	SI, SK, TR, BF,
BJ, CF, CG,	CI, CM, GA,	GN, GQ, GW, ML, MR, NE,	SN, TD, TG, BW,
GH, GM, KE,	LS, MW, MZ,	NA, SD, SL, SZ, TZ, UG,	ZM, ZW, AM, AZ,

BY, KG, KZ, MD, RU, TJ, TM

PRIORITY APPLN. INFO.:

US 2006-782787P P 20060316

AB The present invention relates to thiazolidinedione analogs that are useful for treating hypertension. Thus, pharmaceutical antihypertensive composition comprising a thiazolidinedione analog (15-60 mg), and CM-cellulose, magnesium stearate, hydroxypropyl cellulose, and lactose monohydrate was

formulated.
IT 146062-49-9 950696-94-3 950696-95-4
950696-96-5 950696-97-6 950696-98-7
950696-99-8 950697-00-4 950697-01-5
950697-03-7 950697-05-9 950697-07-1

950697-09-3
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(thiazolidinedione analogs for treating hypertension)

RN 146062-49-9 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-oxoethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

| Et

RN 950696-94-3 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[(2R)-2-(5-ethyl-2-pyridinyl)-2hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 950696-95-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[(2S)-2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 950696-96-5 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-5-methyl- (CA INDEX NAME)

PAGE 1-A

| Et

L8 ANSWER 3 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2007:1090461 CAPLUS

TITLE:

Thiazolidinedione analogues for treating metabolic

inflammation mediated diseases such as diabetes

INVENTOR(S): Colca, Gerard R.; Kletzien, Rolf F.

PATENT ASSIGNEE(S):

Metabolic Solutions Development Company, USA

SOURCE: PCT Int. Appl., 45pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

7

PATENT INFORMATION:

PATENT NO.	KIND DATE				7	APPL:	CAT:		DATE						
WO 20071000	72 20070027			WO 2007-US6321						20070314					
WO 2007109024			A2	•	2007	0,921	,	40 Z	JU / - (J303.	4.1		2.	00,0.	7
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BJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,

GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

PRIORITY APPLN. INFO.:

US 2006-782894P P 20060316

AB The present invention relates to thiazolidinedione analogs that are useful for treating metabolic inflammation mediated diseases such as diabetes. Thus, pharmaceutical composition comprising a thiazolidinedione analog (15-60 mg), and CM-cellulose, magnesium stearate, hydroxypropyl cellulose, and lactose monohydrate was formulated. It exhibited enhanced binding to PPARγ receptors and decreased the glucose, insulin, and triglyceride level in diabetic mice.

IT 146062-49-9 950696-94-3 950696-95-4 950696-96-5 950696-97-6 950696-98-7 950696-99-8 950697-00-4 950697-01-5

950697-03-7 950697-05-9 950697-07-1

950697-09-3

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(thiazolidinedione analogs for treating metabolic inflammation mediated diseases such as diabetes)

RN 146062-49-9 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-oxoethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

| Et

RN 950696-94-3 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[(2R)-2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

Et

ANSWER 4 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2006:1253003 CAPLUS

DOCUMENT NUMBER:

146:804

TITLE:

insulin sensitization for delaying puberty and

increasing growth

INVENTOR (S):

De Zegher, Francis; Dunger, David; Ibanez, Lourdes

K.U. Leuven Research and Development, Belg.;

PATENT ASSIGNEE(S): Addenbrooke's Hospital PCT Int. Appl., 61pp.

SOURCE:

CODEN: PIXXD2 Patent

DOCUMENT TYPE: LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.					KIN	ND DATE			APPLICATION NO.							DATE		
						-												
WO 2006125285					A1		2006	1130	WO 2006-BE60						20060523			
WO 2006125285					B1		2007	0111										
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		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	ΚP,	KR,	
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		MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	
		SG,	SK,	SL,	SM,	SY,	ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UΖ,	VC,	
		VN,	YU,	ZA,	ZM,	zw												

RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

PRIORITY APPLN. INFO.:

GB 2005-10469 A 20050523.

OTHER SOURCE(S): MARPAT 146:804

AB In accordance with the purpose of the invention, as embodied and broadly described herein, the invention is broadly drawn to a new method of treatment, the use of agents to manufacture a composition of treatment or the composition

of treatment for the prevention of rapidly progressive puberty, the prevention of early menarche or the modulation, more particularly the delay, of the tempo of puberty in a female mammal, preferably a human girl, and the disorders related thereto. In a particular embodiment the present invention involves the use of at least one insulin-sensitizing agent such as metformin, any of the polymorphs of metformin or a pharmaceutically acceptable salt thereof for the preparation of a composition

of

treatment to modulate the tempo of pubertal progression in a girl. Metformin administration to girls experiencing precocious puberty resulted in normalization of pubertal progression to menarche, increased height gains, leaner body composition, and decreases indexes relating to insulin resistance.

IT 101931-00-4 105355-33-7 111025-46-8,

Pioglitazone 146062-44-4 146062-45-5

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL

(Biological study); USES (Uses)

(metformin-induced insulin sensitization for delaying puberty and increasing growth)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

Ac

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 5 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN 1.8

6

ACCESSION NUMBER:

2006:799525 CAPLUS

DOCUMENT NUMBER:

145:305561

TITLE:

Pioglitazone is metabolised by CYP2C8 and CYP3A4 in

vitro: potential for interactions with CYP2C8

inhibitors

AUTHOR (S):

Jaakkola, Tiina; Laitila, Jouko; Neuvonen, Pertti J.;

Backman, Janne T.

CORPORATE SOURCE:

Department of Clinical Pharmacology, University of

Helsinki and Helsinki University Central Hospital,

Helsinki, Finland

SOURCE:

Basic & Clinical Pharmacology & Toxicology (2006),

99(1), 44-51

CODEN: BCPTBO; ISSN: 1742-7835

Blackwell Publishing Ltd.

DOCUMENT TYPE:

PUBLISHER:

Journal

LANGUAGE:

English

Our objective was to identify the cytochrome P 450 (CYP) enzymes that metabolize pioglitazone and to examine the effects of the CYP2C8 inhibitors montelukast, zafirlukast, trimethoprim and gemfibrozil on pioglitazone metabolism in vitro. The effect of different CYP isoform inhibitors on the elimination of a clin. relevant concentration of pioglitazone $(1 \mu M)$ and the formation of the main primary metabolite M-IV were studied using pooled human liver microsomes. The metabolism of pioglitazone

by CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, CYP2D6, CYP2E1, CYP3A4 and CYP3A5 was investigated using human recombinant CYP isoforms. In particular, the inhibitors of CYP2C8, but also those of CYP3A4, markedly inhibited the elimination of pioglitazone and the formation of M-IV by HLM. Inhibitors selective to other CYP isoforms had a minor effect only. Of the recombinant isoforms, CYP2C8 (20 pmol/mL) metabolized pioglitazone markedly (56% in 60 min.), and also CYP3A4 had a significant effect (37% in 60 min.). Montelukast, zafirlukast, trimethoprim and gemfibrozil inhibited pioglitazone elimination in HLM with IC50 values of 0.51 μM, 1.0 μM , 99 μM and 98 μM , resp., and the formation of the metabolite M-IV with IC50 values of 0.18 $\mu M,~0.78~\mu M,~71~\mu M$ and 59 μM , resp. In conclusion, pioglitazone is metabolized mainly by CYP2C8 and to a lesser extent by CYP3A4 in vitro. CYP2C9 is not significantly involved in the elimination of pioglitazone. The effect of different CYP2C8 inhibitors on pioglitazone pharmacokinetics needs to be evaluated also in vivo because, irresp. of their in vitro CYP2C8 inhibitory potency, their pharmacokinetic properties may affect the extent of interaction.

IT 176109-96-9

RL: BSU (Biological study, unclassified); BIOL (Biological study) (pioglitazone is metabolized by CYP2C8 and CYP3A4 in vitro: potential for interactions with CYP2C8 inhibitors)

RN 176109-96-9 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-hydroxy-2-[5-(1-hydroxyethyl)-2-pyridinyl]ethoxy]phenyl]methyl]- (9CI) (CA INDEX NAME)

PAGE 1-A

IT 111025-46-8, Pioglitazone

RL: PKT (Pharmacokinetics); BIOL (Biological study)

(pioglitazone is metabolized by CYP2C8 and CYP3A4 in vitro: potential

for interactions with CYP2C8 inhibitors)

RN 111025-46-8 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl](CA INDEX NAME)

PAGE 1-A

PAGE 2-A

Et

REFERENCE COUNT:

THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 6 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2006:370121 CAPLUS

DOCUMENT NUMBER:

145:240842

TITLE:

Simultaneous determination of pioglitazone and its two

active metabolites in human plasma by HPLC-MS

AUTHOR(S):

Deng, Lijing; Wang, Feng; Xie, Zhihong; Xiao, Yiwen;

Li, Huande

CORPORATE SOURCE:

Second Xiangya Hospital, Central-South University, Changsha, Hunan Province, 410011, Peop. Rep. China Zhongguo Yaoxue Zazhi (Beijing, China) (2005), 40(10),

SOURCE:

772-774

CODEN: ZYZAEU; ISSN: 1001-2494

PUBLISHER:

Zhongguo Yaoxue Zazhishe

DOCUMENT TYPE:

Journal Chinese

LANGUAGE:

10/520,166

A HPLC-MS method was established for simultaneous determining of pioglitazone and its two active metabolites: M-III (keto-derivative) and M-IV (hydroxy-derivative) in human plasma. The separation was performed on a Waters XTerraTM C18 column (2.1 mm+150 mm, 3.5 μm) with guard column Phenomenex C18. The column temperature was 50 degree. The mobile phase consisted of acetonitrile-30 mmol·L-1 ammonium acetate solution (added with 0.1% formic acid and 0.05% trifluoroacetic acid) (35:65), with a flow rate of 0.22 mL·min-1. The compound was ionized in the electrospray ionization (ESI) ion source of the mass spectrometer and selected ion mass spectral (m/z) 357.4 (PIO), 358.2 (is), 371.5 (M-III), 373.2 (M-IV) to quantify. Human plasma samples were extracted with 1:2 chlorform:methyl t-Bu ether after acidification. The linear ranges were 11.16-1748.60 ng·mL-1 for pioglitazone, 3.33-520.50 ng·mL-1 for M-III and 5.00-687.50 ng·mL-1 for M-IV (r≥0.9997), and their detect limits were 2.90, 1.10, 1.20 ng mL-1. Recoveries were within 90%-110%, and intra-and inter-day RSDs were all less than 15%. The method is found to be sensitive, rapid and accurate, and has been applied successfully to sample anal. for clin. study of pioglitazone pharmacokinetics and drug interaction.

IT 101931-00-4 146062-45-5

RL: PKT (Pharmacokinetics); BIOL (Biological study) (simultaneous determination of pioglitazone and its two active metabolites

in

human plasma by HPLC-MS)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

RN 146062-45-5 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-acetyl-2-pyridinyl)ethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

I Ac

IT 111025-46-8, Pioglitazone

RL: PKT (Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(simultaneous determination of pioglitazone and its two active metabolites

in

human plasma by HPLC-MS)

RN 111025-46-8 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-(CA INDEX NAME)

Et

L8 ANSWER 7 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2006:72865 CAPLUS

DOCUMENT NUMBER:

145:20356

TITLE:

Effect of rifampicin on the pharmacokinetics of

pioglitazone

AUTHOR(S):

Jaakkola, Tiina; Backman, Janne T.; Neuvonen, Mikko;

Laitila, Jouko; Neuvonen, Pertti J.

CORPORATE SOURCE:

Department of Clinical Pharmacology, University of Helsinki and Helsinki University Central Hospital,

Helsinki, Finland

SOURCE:

British Journal of Clinical Pharmacology (2006),

61(1), 70-78

CODEN: BCPHBM; ISSN: 0306-5251

PUBLISHER:

Blackwell Publishing Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB Aims The effect of enzyme induction on the pharmacokinetics of pioglitazone, a thiazolidinedione antidiabetic drug that is metabolized primarily by CYP2C8, is not known. Rifampicin is a potent inducer of several CYP enzymes and our objective was to study its effects on the pharmacokinetics of pioglitazone in humans. Methods In a randomized, two-phase crossover study, ten healthy subjects ingested either 600 mg rifampicin or placebo once daily for 6 days. On the last day, they received a single oral dose of 30 mg pioglitazone. The plasma concns. and cumulative excretion of pioglitazone and its active metabolites M-IV and M-III into urine were measured up to 48 h. Results Rifampicin decreased

the mean total area under the plasma concentration-time curve (AUC0-∞) of pioglitazone by 54% (range 20-66%; P = 0.0007; 95% confidence interval -78 to -30%) and shortened its dominant elimination half-life (t1/2) from 4.9 to 2.3 h (P = 0.0002). No significant effect on peak concentration (Cmax) or time to peak (tmax) was observed Rifampicin increased the apparent formation rate of M-IV and shortened its tmax (P < 0.01). It also decreased the $AUC0-\infty$ of M-IV (by 34%; P = 0.0055) and M-III (by 39%; P = 0.0026), shortened their t1/2 (M-IV by 50%; P = 0.0008, and M-III by 55%; P = 0.0016) and increased the AUCO-∞ ratios of M-IV and M-III to pioglitazone by 44% (P = 0.0011) and 32% (P = 0.0027), resp. Rifampicin increased the M-IV/pioglitazone and M-III/pioglitazone ratios in urine by 98% (P = 0.0015) and 95% (P = 0.0024). A previously unrecognized metabolite M-XI, tentatively identified as a dihydroxy metabolite, was detected in urine during both phases, and rifampicin increased the ratio of M-XI to pioglitazone by 240% (P = 0.0020). Conclusions Rifampicin caused a substantial decrease in the plasma concentration of pioglitazone, probably by induction of CYP2C8. Concomitant use of rifampicin with pioglitazone may decrease the efficacy of the latter drug.

IT 111025-46-8, Pioglitazone

RL: PAC (Pharmacological activity); PKT (Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (rifampicin decreased AUC, t1/2 of pioglitazone and its metabolites

M-III, M-IV, did not affect Cmax, tmax of pioglitazone, increased Kf of M-IV, increased M-IV/pioglitazone, M-III/pioglitazone ratios in healthy human)

RN 111025-46-8 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl](CA INDEX NAME)

PAGE 1-A

CH-- Me OH

REFERENCE COUNT:

THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 8 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

30

ACCESSION NUMBER:

2005:1259919 CAPLUS

DOCUMENT NUMBER:

144:324098

TITLE:

Effect of gemfibrozil on the pharmacokinetics of

pioglitazone

AUTHOR(S):

Deng, Li-Jing; Wang, Feng; Li, Huan-De

CORPORATE SOURCE:

Clinical Pharmaceutical Research Institute, The Second

Xiangya Hospital, The Central South University,

Changsha, 410011, Peop. Rep. China

SOURCE:

European Journal of Clinical Pharmacology (2005),

61(11), 831-836

CODEN: EJCPAS; ISSN: 0031-6970

PUBLISHER:

Springer Journal

DOCUMENT TYPE: LANGUAGE: English

Objective: Our objective was to study the effects of gemfibrozil on the pharmacokinetics of pioglitazone and the active compds., which are all the substrates of CYP2C8 and CYP3A4. Methods: In a randomized, two-phase crossover study, 10 healthy volunteers were pretreated for 2 days with either 600 mg oral gemfibrozil or placebo twice daily. On day 3, they received a single dose of 600 mg gemfibrozil or placebo, and 1 h later they received a single oral dose of 30 mg pioglitazone. Plasma concns. of

10/520,166

pioglitazone and both active metabolites M-III and M-IV were measured for up to 120 h. Results: Gemfibrozil raised the mean total area under the concentration-time curve (AUC) of parent pioglitazone 3.4-fold (P<0.001). No statistically significant changes were seen in the total AUC of M-III or M-IV after gemfibrozil pretreatment. Gemfibrozil reduced the M-III/pioglitazone and M-IV/pioglitazone AUC0- ∞ ratio by 71% (P<0.001) and 65% (P<0.001), strikingly prolonging their t1/2. Conclusion: Gemfibrozil greatly increased the plasma concentration of parent pioglitazone

and

also inhibited the further metabolism of M-III and M-IV. Careful blood glucose monitoring and dosage adjustments are suggested during coadministration of pioglitazone and gemfibrozil.

IT 101931-00-4 146062-45-5

RL: BSU (Biological study, unclassified); PKT (Pharmacokinetics); BIOL (Biological study)

(gemfibrozil inhibited metabolism of pioglitazone metabolite M-III and M-IV showing that careful blood glucose monitoring and dosage adjustment are required during their coadministration in human)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

Et.

Ac

111025-46-8, Pioglitazone IT

RL: PKT (Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(gemfibrozil markedly increased pioglitazone plasma concentration while inhibited M-III and M-IV metabolism showing that careful blood glucose monitoring and dosage adjustment are required during their coadministration in human)

RN 111025-46-8 CAPLUS

2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-CN(CA INDEX NAME)

Et

REFERENCE COUNT:

THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS 15 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 9 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:1139098 CAPLUS

DOCUMENT NUMBER:

144:141941

TITLE:

Single- and multiple-dose pharmacokinetics of pioglitazone in adolescents with type 2 diabetes Christensen, Michael L.; Meibohm, Bernd; Capparelli,

AUTHOR (S):

Edmund V.; Velasquez-Mieyer, Pedro; Burghen, George

A.; Tamborlane, William V.

CORPORATE SOURCE:

Pediatric Pharmacology Research Units, The University of Tennessee and LeBonheur Children's Medical Center,

Memphis, USA

SOURCE:

Journal of Clinical Pharmacology (2005), 45(10),

1137-1144

CODEN: JCPCBR; ISSN: 0091-2700

PUBLISHER:

Sage Publications

DOCUMENT TYPE:

Journal English

LANGUAGE:

This study assessed the single- and multiple-dose pharmacokinetics of 3 doses (15 mg, 30 mg, and 45 mg) of pioglitazone in 36 adolescents with type 2 diabetes. Blood samples were obtained over a 48-h interval after the first dose (day 1) and over a 72-h interval after the last dose (day

15) of pioglitazone and were assayed for pioglitazone and active metabolites (M-III and M-IV). Pioglitazone systemic exposure increased dose dependency but was less than dose proportional during multiple

10/520,166

dosing. The median peak pioglitazone concentration occurred at 2 h. The mean half-life was 8 to 9 h for pioglitazone and 24 to 32 h for M-III and M-IV, with similar values at each dose level. During multiple dosing, accumulation for pioglitazone was negligible, but it reached 2.5- to 3.0-fold for M-III and M-IV. The sustained total serum concentration of active compds. during multiple dosing provides the basis for once-daily dose administration of pioglitazone in adolescents.

IT 146062-45-5

RL: BSU (Biological study, unclassified); PKT (Pharmacokinetics); BIOL (Biological study)

(pharmacokinetic anal. of pioglitazone metabolite M-III revealed less dose-proportional increase in Cmax and AUC during multiple dosing compared to single dosing in adolescent with type 2 diabetes)

RN 146062-45-5 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-acetyl-2-pyridinyl)ethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

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IT 101931-00-4

RL: BSU (Biological study, unclassified); PKT (Pharmacokinetics); BIOL (Biological study)

(pharmacokinetic anal. of pioglitazone metabolite M-IV revealed less dose-proportional increase in Cmax and AUC during multiple dosing compared to single dosing in adolescent with type 2 diabetes)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

| Et

(CA INDEX NAME)

Et

REFERENCE COUNT:

22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 10 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:60504 CAPLUS

DOCUMENT NUMBER:

140:111409

TITLE:

A novel process to prepare pioglitazone via several

novel intermediates.

INVENTOR(S):

Pandey, Bipin; Lohray, Vidya Bhushan; Lohray, Braj

Bhushan

PATENT ASSIGNEE(S):

Cadila Healthcare Limited, India

PCT Int. Appl., 74 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.					KIN	D	DATE		APPLICATION NO.							DATE			
	2004				A2 A3		2004 2004		WO 2003-IN241						20030715				
WO		ΑE,	AG,		AM,	AT,		AZ,	-	-									
		GM,	HR,	HU,	ID,	IL,	IN, MD,	IS,	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,		
		PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	TJ,	TM,	TN,		

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TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
              KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
              FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
              BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                                IN 2002-MU648
                                                                          20020716
     IN 2002MU00648
                            Α
                                   20040424
     AU 2003272072
                            A1
                                   20040202
                                               . AU 2003-272072
                                                                          20030715
                                 20050413
     EP 1521753
                            A2
                                                EP 2003-753913
                                                                          20030715
     EP 1521753
                            B1
                                   20070905
              AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
                            Т
                                   20070915
                                                AT 2003-753913
                                                                          20030715
     AT 372336
                                   20060727
                                                US 2005-520166
                                                                          20051004
     US 2006167061
                            A1
PRIORITY APPLN. INFO.:
                                                IN 2002-MU648
                                                                      A 20020716
                                                WO 2003-IN241
                                                                      W 20030715
                           CASREACT 140:111409; MARPAT 140:111409
OTHER SOURCE(S):
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- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- AB The present invention discloses a novel and general process to prepare various pyridine substituted 5-[4-[2-(alkyl substituted pyridyl)ethoxy]benzyl]-2,4-thiazolidinone derivs. of general formula I [R = alkyl], and their pharmaceutically acceptable salts. The present invention especially provides a novel process to prepare pioglitazone hydrochloride [R = 5-ethyl], via novel intermediates, i.e. II and III. This process involves lesser number of steps with high yields and uses key solid intermediates, which are operationally simple, and therefore offers opportunities for better com. viability.
- IT 101931-00-4P 111025-46-8P 646519-87-1P 646519-88-2P 646519-89-3P 646519-91-7P 646519-93-9P
 - RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of pioglitazone via several novel intermediates)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

L8 ANSWER 11 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003:766616 CAPLUS

DOCUMENT NUMBER: 140:192140

TITLE: Sex differences in the pharmacokinetics of

Pioglitazone in rats

AUTHOR(S): Fujita, Yukiyoshi; Yamada, Yasuhiko; Kusama, Makiko;

Yamauchi, Toshimasa; Kamon, Junji; Kadowaki, Takashi;

Iga, Tatsuji

CORPORATE SOURCE: Faculty of Medicine, University of Tokyo Hospital,

Department of Pharmacy, University of Tokyo,

Bunkyo-ku, Tokyo, Japan

SOURCE: Comparative Biochemistry and Physiology, Part C:

Toxicology & Pharmacology (2003), 136C(1), 85-94

CODEN: CBPPFK; ISSN: 1532-0456

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

AB Clin. studies have suggested that Pioglitazone, an insulin sensitizer, has a stronger effect in women than in men. To determine the sex difference in the pharmacokinetics of Pioglitazone, the authors examined the plasma and white adipose tissue levels of Pioglitazone and its active metabolites (M-II, M-III, and M-IV) in male and female rats treated with a single or repeated oral administration of Pioglitazone (10 mg/kg). The AUCs of Pioglitazone $(149.6\pm22.6 \text{ vs. } 103.3\pm14.0 \text{ } \mu\text{g}\cdot\text{h/mL}; \text{ P<0.01}), \text{ M-III}$ $(31.4\pm8.1 \text{ vs. } 20.2\pm4.7 \text{ } \mu\text{g}\cdot\text{h/mL}; \text{ P<0.05}), \text{ and M-IV}$ $(41.9\pm15.5 \text{ vs. } 14.1\pm1.6 \text{ } \mu\text{g}\cdot\text{h/mL}; \text{ P<0.01})$ were larger in female rats than in male rats, but the levels of M-II were similar. Any of the compds. did not accumulate in plasma after repeated administration. According to kinetic model anal., the apparent elimination rate of Pioglitazone and the formation rate of M-II were faster in male rats than in female rats. No significant sex difference was found in the tissue-to-plasma concentration ratios of Pioglitazone or its active metabolites in white adipose tissue. These results suggest that there are sex differences in the plasma levels of Pioglitazone and some of its active metabolites and that those differences are reflected in differences in white adipose tissue levels.

IT 101931-00-4 111025-46-8D, Pioglitazone, metabolites 146062-44-4 146062-45-5

RL: PKT (Pharmacokinetics); BIOL (Biological study)
(Pioglitazone and its active metabolites in blood plasma and white
adipose tissue of male and female rats after oral administration of
Pioglitazone)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

Et

RN 111025-46-8 CAPLUS CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-(CA INDEX NAME)

Εt

REFERENCE COUNT:

CORPORATE SOURCE:

SOURCE:

THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2007 ACS on STN ANSWER 12 OF 26

33

2003:389760 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 140:238

Identification of novel metabolites of pioglitazone in TITLE:

rat and dog

Shen, Z.; Reed, J. R.; Creighton, M.; Liu, D. Q.; AUTHOR (S):

Tang, Y. S.; Hora, D. F.; Feeney, W.; Szewczyk, J.;

Bakhtiar, R.; Franklin, R. B.; Vincent, S. H. Departments of Drug Metabolism, Merck Research

Laboratories, Rahway, NJ, 07065, USA

Xenobiotica (2003), 33(5), 499-509

CODEN: XENOBH; ISSN: 0049-8254

PUBLISHER: Taylor & Francis Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

Four new metabolites of pioglitazone were identified by liquid AB chromatog.-mass spectrometry (LC-MS/MS) as being formed by hydroxylation (M-VII and M-VIII), opening of the thiazolidinedione ring (M-X) and by desatn. of the terminal Et side chain or tether ethoxy moiety (M-IX), resp. The structure of one of the hydroxylated metabolites (M-VII) was confirmed by chemical modification using the Jones reaction. Oxidative cleavage of the thiazolidinedione ring is a novel pathway not previously reported for pioglitazone. The hydroxylated M-VII was detected in incubations with rat, dog and human liver and kidney microsomes, and in

10/520,166

plasma from rats and dogs dosed orally with [3H]pioglitazone. The carboxylic acid derivative of M-VII (M-V) and its taurine conjugate were the major radioactive components in dog bile.

IT 101931-00-4 146062-44-4 146062-45-5

146062-48-8 157142-91-1 186751-40-6

625853-75-0 625853-76-1

RL: BSU (Biological study, unclassified); BIOL (Biological study) (Identification and localization of novel metabolites of pioglitazone in rat and dog)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

| Et

RN 146062-44-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-[5-(1-hydroxyethyl)-2-pyridinyl]ethoxy]phenyl]methyl]- (CA INDEX NAME)

Εt

REFERENCE COUNT:

19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2007 ACS on STN L8ANSWER 13 OF 26

ACCESSION NUMBER:

2002:610859 CAPLUS

DOCUMENT NUMBER:

137:312674

TITLE:

AUTHOR(S):

Process Development and Scale-Up of the Potential Thiazolidinedione Antidiabetic Candidate PNU-91325 Carpenter, Donald E.; Imbordino, Rick J.; Maloney, Mark T.; Moeslein, Jeffery A.; Reeder, Michael R.;

Scott, Allen

CORPORATE SOURCE:

Early Process Research and Development, Pharmacia

Corporation, Kalamazoo, MI, 49001, USA

SOURCE:

Organic Process Research & Development (2002), 6(5),

721-728

CODEN: OPRDFK; ISSN: 1083-6160

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

English

LANGUAGE:

An efficient six-step synthesis was developed for the preparation of the thiazolidinedione analog PNU-91325 from a com. available olefin. This process involves a novel epoxide ring opening with a deactivated phenol under phase-transfer conditions. Significant improvements were made in the oxidation of a secondary alc. to the ketone and the 1,4-reduction of an enone

from a previous process. Overall, this route allows for the preparation of PNU-91325 in 25% yield.

IT 146062-49-9P

RL: IMF (Industrial manufacture); PREP (Preparation) (potential antidiabetic agent; process development and scale-up of potential thiazolidinedione antidiabetic candidate PNU-91325)

RN 146062-49-9 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-oxoethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

| Et

IT 101931-00-4P

RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)

(process development and scale-up of potential thiazolidinedione antidiabetic candidate PNU-91325)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

Et

REFERENCE COUNT:

THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS 21 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 14 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

1

ACCESSION NUMBER:

1999:753074 CAPLUS

DOCUMENT NUMBER:

131:346538

TITLE:

Thiazolidine and oxazolidine derivatives for the

treatment of acute myocardial infarction and

inhibition of cardiomyocyte apoptosis

INVENTOR(S):

Wang, Ping H.

PATENT ASSIGNEE(S): SOURCE:

Regents of the University of California, USA

PCT Int. Appl., 49 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.					KIND DATE					APPLICATION NO.						DATE		
WO 9959586				Al		1999	1125	WO 1999-US11101						19990519				
	W:	AE,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	
		DE,	DK,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	
		JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	
		MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	
		TM,	TR,	TT,	UA,	UG,	US,	UZ,	VN,	ΥU,	ZA,	ZW,	AM,	ΑZ,	BY,	KG,	ΚZ,	
		MD,	RU,	TJ,	TM													

RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,

CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

AU 9940052 A 19991206 AU 1999-40052 19990519
PRIORITY APPLN. INFO.: US 1998-86030P P 19980519
US 1998-87204P P 19980528
WO 1999-US11101 W 19990519

OTHER SOURCE(S): MARPAT 131:346538

AB It has been demonstrated that antidiabetic thiazolidine and oxazolidine derivs. (glitazones) exhibit novel effects on apoptosis of cardiomyocytes. These substances are capable of greatly decreasing apoptosis by a pathway that is not Caspase 3 dependent. Addition of IGF1 to the treatment further prevents apoptosis. Glitazones alone or glitazones plus IGF1 should be administered at the beginning of a myocardial infarction and continued through the recuperation period to reduce morbidity and prevent unfavorable remodeling of the myocardium. Thus, troglitazone (5 $\mu \rm M)$, when added to a culture medium, reduced doxorubicin-induced apoptosis of cardiomyocyte by approx. 60%.

IT 74773-17-4 101930-98-7 101931-00-4

111025-46-8, Pioglitazone

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(thiazolidine and oxazolidine derivs. for treatment of acute myocardial infarction and inhibition of cardiomyocyte apoptosis)

RN 74773-17-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(2-pyridinyl)ethoxy]phenyl]methyl]- (9CI) (CA INDEX NAME)

CN

RN 101930-98-7 CAPLUS

2,4-Thiazolidinedione, 5-[[4-[2-hydroxy-2-(6-methyl-2-pyridinyl)ethoxy]phenyl]methyl]- (9CI) (CA INDEX NAME)

RN 101931-00-4 CAPLUS
CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

Et

RN 111025-46-8 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-(CA INDEX NAME)

PAGE 1-A

PAGE 2-A

| Et

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 15 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:485977 CAPLUS

DOCUMENT NUMBER:

131:252097

TITLE:

Three-dimensional quantitative structure activity relationships (3-D-QSAR) of antihyperglycemic agents Kulkarni, Santosh S.; Gediya, Lalji K.; Kulkarni,

AUTHOR(S):

Vithal M.

CORPORATE SOURCE:

Pharmaceutical Division, Department of Chemical Technology, University of Mumbai, Mumbai, 400 019,

India

SOURCE:

Bioorganic & Medicinal Chemistry (1999), 7(7),

1475-1485

CODEN: BMECEP; ISSN: 0968-0896

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

A three-dimensional quant. structure activity relationship study (3-D-QSAR) was performed on a set of thiazolidinedione antihyperglycemic agents using the comparative mol. field anal. (COMFA) method. The COMFA models were derived from a training set of 53 compds. Fifteen compds., which were not used in model generation were used to validate the COMFA models. All the compds. were superimposed to the template structure by atom-based and shape-based strategies. The SYBYL QSAR rigid body field fit was also used for aligning the ligands. A total of twelve different alignments were generated. The resulting models exhibited a good cross-validated rcv2 values (0.624-0.764) and the conventional r2 values (0.689-0.921). A more robust cross-validation test using cross-validation by 2 groups (leave half out method) was performed 100 times to ascertain the predictiveness of the CoMFA models. The mean of rcv2 values from 100 runs ranged from 0.611-0.690. Few models exhibited good external predictivity. These models were then used to define a hypothetical receptor model for antihyperglycemic agents.

IT 74773-17-4 74773-19-6 101930-98-7

101930-99-8 101931-00-4 101931-04-8

101931-05-9 101946-34-3 101946-35-4

105355-33-7 105355-34-8 105355-35-9

111025-46-8 127676-13-5 127676-14-6

127676-15-7

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(3-D-QSAR of antihyperglycemic agents)

RN 74773-17-4 CAPLUS

2,4-Thiazolidinedione, 5-[[4-[2-(2-pyridinyl)ethoxy]phenyl]methyl]- (9CI) (CA INDEX NAME)

CN

CN

RN 74773-19-6 CAPLUS

2,4-Thiazolidinedione, 5-[[4-[2-(6-methyl-2-pyridinyl)ethoxy]phenyl]methyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 16 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1997:93456 CAPLUS

DOCUMENT NUMBER:

126:287573

TITLE:

Disposition of the new antidiabetic agent pioglitazone

in rats, dogs, and monkeys

AUTHOR(S):

Maeshiba, Yoshihiro; Kiyota, Yutaka; Yamashita, Kenji;

Yoshimura, Yoshinobu; Motohashi, Michio; Tanayama,

Shigeharu

CORPORATE SOURCE:

Drug Analysis Pharmacokinetics Research Laboratories, Takeda Chemical Industries Ltd., Osaka, 532, Japan

SOURCE:

Arzneimittel-Fórschung (1997), 47(1), 29-35

CODEN: ARZNAD; ISSN: 0004-4172

PUBLISHER:
DOCUMENT TYPE:

LANGUAGE:

to

Cantor Journal English

The disposition of pioglitazone was studied after oral administration to rats, dogs, and monkeys using 14C-labeled drug. After oral dosing, pioglitazone was well absorbed from the gastrointestinal tract at an extent of 96, 95, and 90% in rats, dogs, and monkeys, resp. In rats, the concentration of pioglitazone in plasma reached a peak (Cmax 0.71 μg/mL) at 4 h (tmax) after dosing and declined with a half-life (t1/2) of 2.6 h. In dogs, tmax, Cmax and t1/2 were 0.5 h, 0.32 μg/mL and 2.1 h, and those for monkeys were 4.3 h, 0.48 μg/mL and 5.3 h, resp. The drug was metabolized mainly to M-I to M-VI including the pharmacol. active metabolites (M-II, III, and IV). The pharmacol. active compds. (total of the unchanged compound and the above three active metabolites) accounted for 87, 71 and 73% of the radioactivity in plasma of rats, dogs, and monkeys, resp. The radioactivity was widely distributed in tissues after oral administration to rats, and decreased to the very low concentration within 24

72 h after dosing. Radioactivity dose was almost completely excreted in urine and feces.

IT 101931-00-4 146062-44-4 146062-45-5
RL: BPR (Biological process); BSU (Biological study, unclassified); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation,

nonpreparative); PROC (Process)

(disposition of antidiabetic pioglitazone in rats, dogs, and monkeys)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

Et

RN 146062-44-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-[5-(1-hydroxyethyl)-2-pyridinyl]ethoxy]phenyl]methyl]- (CA INDEX NAME)

CO₂H

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CAPLUS COPYRIGHT 2007 ACS on STN
    ANSWER 17 OF 26
L8
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ACCESSION NUMBER: 1997:93455

DOCUMENT NUMBER:

126:287572

TITLE:

Studies on the metabolism of the new antidiabetic agent pioglitazone. Identification of metabolites in

rats and dogs

AUTHOR(S):

Kiyota, Yutaka; Kondo, Takahiro; Maeshiba, Yoshihiro; Hashimoto, Ai; Yamashita, Kenji; Yoshimura, Yoshinobu;

Motohashi, Michio; Tanayama, Shigeharu

CORPORATE SOURCE:

Drug Analysis Pharmacokinetics Research Laboratories, Takeda Chemical Industries Ltd., Osaka, 532, Japan

Arzneimittel-Forschung (1997), 47(1), 22-28

CODEN: ARZNAD; ISSN: 0004-4172

CAPLUS

SOURCE:

Cantor

PUBLISHER: DOCUMENT TYPE:

Journal LANGUAGE: English

Metabolic studies of pioglitazone,, a new antidiabetic agent, in rats and dogs using liquid chromatog./tandem mass spectrometry and 1H-NMR led to characterization of the following metabolites: the parent compound, (\pm) -5-(p-hydroxybenzyl)-2,4-thiazolidinedione (M-I), (\pm) -5-[p-[2-(5-ethyl-2-pyridyl)-2-hydroxyethoxy]benzyl]-2,4-

thiazolidinedione (M-II), (\pm) -5-[p-[2-(5-acetyl-2pyridyl) ethoxy] benzyl] -2,4-thiazolidinedione (M-III), (\pm) -5-[p-[2-[5-(1-pyridyl)]] hydroxyethyl)-2-pyridyl]ethoxy]benzyl]-2,4-thiazolidinedione (M-IV),

 (\pm) -5-[p-[2-(5-carboxymethyl-2-pyridyl)ethoxy]-benzyl]-2,4thiazolidinedione (M-V), and (\pm) -5-[p-[2-(5-carboxy-2IT

pyridyl)ethoxy]benzyl]-2,4-thiazolidinedione (M-VI). Pioglitazone is considered to be metabolized by cleavage of aliphatic CO bond to lead to M-I, hydroxylation of aliphatic methylene groups to form M-II and M-IV, oxidation of M-IV to give M-III, oxidation of the Et group to form M-V, and oxidative loss of the terminal carbon to lead to M-VI. Furthermore, part of metabolites exist as conjugated form. Among the conjugates, M-IV conjugated with sulfuric acid and M-V conjugated with taurine were identified.

111025-46-8, Pioglitazone RL: BPR (Biological process); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (metabolism of antidiabetic pioglitazone and identification of metabolites in rats and dogs)

111025-46-8 CAPLUS RN

2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-CN (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

Εt

101931-00-4 146062-44-4 146062-45-5 IT

146062-48-8 186751-40-6

RL: BSU (Biological study, unclassified); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative)

(metabolism of antidiabetic pioglitazone and identification of metabolites in rats and dogs)

101931-00-4 CAPLUS RN CN

2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2hydroxyethoxy]phenyl]methyl] - (CA INDEX NAME)

CO₂H

ANSWER 18 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1997:57026 CAPLUS

DOCUMENT NUMBER:

126:152372

TITLE: .

Disposition of AD-4833, a new antidiabetic agent, in

animals

AUTHOR (S):

Maeshiba, Yoshihiro; Kiyota, Yutaka; Yamashita, Kenji;

Yoshimura, Yoshinobu; Motohashi, Michio; Tanayama,

Shigeharu

CORPORATE SOURCE:

Drug analysis and Pharmacokinetics Research

Laboratories, Takeda Chemical Industries, Ltd., Japan

Yakuri to Chiryo (1996), 24(12), 2597-2617

CODEN: YACHDS; ISSN: 0386-3603

PUBLISHER: DOCUMENT TYPE: Raifu Saiensu Shuppan K.K.

Journal

LANGUAGE:

SOURCE:

Japanese

GI

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

The disposition of AD-4833 (I) was studied after oral administration to AB mice, rats, dogs, and monkeys, using [14C] AD-4833-HCl. Concns. of AD-4833 in mouse plasma attained a peak (Cmax, 0.51 $\mu g/mL$) 1.0 h (Tmax) after administration and declined with an apparent half-life (t1/2) of 3.1 h. Concns. of AD-4833 in rat plasma attained a peak (Cmax, 0.71 μ g/mL) 4.0 h (Tmax) after administration and declined with t1/2 of 2.6 h. Tmax, Cmax, and t1/2 in dogs were 0.5 h, 0.32 μ g/mL, and 2.1 h, resp., and those in monkeys were 4.3 h, 0.48 $\mu g/mL$, and 5.3 h, resp. The bioavailabilities of AD-4833 in mice, rats, dogs, and monkeys were 81, 85, 94, and 81%, resp. The pharmacokinetics of AD-4833 in rats and dogs were linear in the dose ranges of 0.5-30 mg/kg and 0.1-3 mg/kg, resp. In rats, the radioactivity was distributed widely in tissues. The concns. of radioactivity in most tissues were lower than those in plasma. The main component in the tissues was the unchanged drug. During repeated administration of [14C]AD-483-HCl for 14 days to rats, the radioactivities in most tissues increased to attain a steady state within the last administration period. After ceasing the administration, 14C levels in tissues declined gradually. The binding of AD-4833 to plasma protein in mice, rats, dogs, and monkeys, and to serum protein in humans, was >98%. The drug was metabolized mainly to 6 compds., three of which were pharmacol. active. The main component in plasma was unchanged AD-4833 in mice, rats, and monkeys, and a metabolite in dogs. Only small amts. of unchanged drug were excreted in the urine and feces of mice, rats, dogs, and monkeys. The excretion of radioactivity in urine and feces was almost complete within 72, 72, 96, and 168 h in mice, rats, dogs, and monkeys, resp. Biliary excretion and enterohepatic circulation were observed in rats. In rats, the radioactivity was excreted quant. within 48 h after the end of repeated administration of [14C] AD-4833-HCl/day for 7 days. 14C was also detected in the milk of rats.

IT 112529-15-4, AD 4833

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(pharmacokinetics and metabolism of)

RN112529-15-4 CAPLUS

2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-CN , hydrochloride (1:1) (CA INDEX NAME)

HCl

101931-00-4 146062-44-4 146062-45-5 TT

146062-48-8 186751-40-6

RL: BPR (Biological process); BSU (Biological study, unclassified); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(pharmacokinetics and metabolism of antidiabetic AD 4833 in relation to formation of)

101931-00-4 CAPLUS RN

2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-CN hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

CO₂H

CAPLUS COPYRIGHT 2007 ACS on STN ANSWER 19 OF 26 L8

ACCESSION NUMBER:

CORPORATE SOURCE:

1996:712946 CAPLUS

DOCUMENT NUMBER:

126:42525

TITLE:

Synthesis and Biological Activity of Metabolites of the Antidiabetic, Antihyperglycemic Agent Pioglitazone Tanis, Steven P.; Parker, Timothy T.; Colca, Jerry R.;

AUTHOR(S):

Fisher, Roberta M.; Kletzein, Rolf F.

SOURCE:

Department of Discovery Chemistry, Pharmacia and

Upjohn Inc., Kalamazoo, MI, 49001, USA Journal of Medicinal Chemistry (1996), 39(26),

5053-5063

PUBLISHER:

CODEN: JMCMAR; ISSN: 0022-2623 American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 126:42525

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- The authors have developed improved syntheses of pioglitazone metabolites AΒ I, II, III, and IV and the putative metabolite ketone V. These entities have been compared in the KKAy mouse model of human type-II diabetes to

CN

pioglitazone. Ketone V has proven to be the most potent of these thiazolidinediones in this in vivo assay. When I-V were compared in vitro in the 3T3-L1 cell line to pioglitazone for their ability to augment insulin-stimulated lipogenesis, V was again the most potent compound with I, II, and IV roughly equivalent to pioglitazone. These data suggest that metabolites I, II, and IV are likely to contribute to the pharmacol. activity of pioglitazone, as had been previously reported for ciglitazone.

IT 101931-00-4P 111025-46-8DP, Pioglitazone, derivs.

146062-44-4P 146062-45-5P 146062-48-8P

146062-49-9P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(synthesis and biol. activity of metabolites of antidiabetic antihyperglycemic agent pioglitazone)

RN 101931-00-4 CAPLUS

2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

| Et

NC-CH2

REFERENCE COUNT:

THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 20 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN L8

37

ACCESSION NUMBER:

1996:320215 CAPLUS

DOCUMENT NUMBER:

125:48213

TITLE:

Simultaneous quantitation of pioglitazone and its metabolites in human serum by liquid chromatography

and solid phase extraction Zhong, W. Z.; Williams, M. G.

CORPORATE SOURCE:

Drug Metabolism Research, Pharmacia & Upjohn, Inc.,

Kalamazoo, MI, 49001, USA

SOURCE:

AUTHOR(S):

Journal of Pharmaceutical and Biomedical Analysis

(1996), 14(4), 465-473

CODEN: JPBADA; ISSN: 0731-7085

PUBLISHER:

Elsevier Journal

English

DOCUMENT TYPE: LANGUAGE:

A high-performance liquid chromatog. (HPLC) method for the simultaneous determination of pioglitazone (U-72107) and its potential metabolites (M-1 to M-6)

in human serum was developed. The method involved a solid phase extraction (SPE) of pioglitazone, its metabolites, and the internal standard (U-92573) from serum using C18 SPE columns with an elution solvent of 0.5 mL of acetonitrile-water (35:65, volume/volume). Separation of the eight analytes

was

AB

achieved within 20 min using a reversed-phase Zorbax RC-C8 anal. column

(250 mm + 4.6 mm i.d., 5 μ m particle size) with a mobile phase of acetonitrile-water (40:60, volume/volume) containing 3 mL acetic acid per L mobile

phase (apparent pH 5.5). An UV detector operated at 269 nm was used with a linear response observed from 0.02 to 2 μg mL-1 for these analytes except for M-4 which was best fitted with a polynomial regression. Limit of quantitation was 0.02 μg mL-1 for pioglitazone, M-3, M-5, and M-6; 0.04 μg mL-1 for M-2 and M-4; and 0.5 μg mL-1 for M-1 when using a 0.5 mL serum sample for extraction Obtained from the method validation, intra-and inter-assay precision was $\leq 9\%$ and accuracy ranged from -8.2 to 13.4% for all analytes. The applicability of this method has been demonstrated by successfully analyzing clin. serum samples. The strategies in the HPLC characterization and in the SPE procedure development for this method are discussed as well.

IT 111025-46-8, Pioglitazone

RL: ANT (Analyte); ANST (Analytical study)

(simultaneous quantitation of pioglitazone and its metabolites in human serum by liquid chromatog. and solid phase extraction)

RN 111025-46-8 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-(CA INDEX NAME)

PAGE 1-A

PAGE 2-A

| Et

IT 101931-00-4 146062-44-4 146062-45-5 146062-48-8 146062-49-9

RL: ANT (Analyte); BSU (Biological study, unclassified); MFM (Metabolic formation); ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative)

| Et

L8 ANSWER 21 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

.1996:215289 CAPLUS

DOCUMENT NUMBER:

124:306331

TITLE:

High-performance liquid chromatographic determination of pioglitazone and its metabolites in human serum and

urine

AUTHOR (S):

Yamashita, Kenji; Murakami, Hiromi; Okuda, Teruaki;

Motohashi, Michio

CORPORATE SOURCE:

Drug Analysis Pharmacokinetics Research Laboratories, Takeda Chemical Industries, Ltd., Osaka, 532, Japan Journal of Chromatography, B: Biomedical Applications

SOURCE:

and

(1996), 677(1), 141-6

PUBLISHER:

CODEN: JCBBEP; ISSN: 0378-4347 Elsevier Journal

DOCUMENT TYPE: LANGUAGE:

NGUAGE: English
A high-performance liquid chromatog. (HPLC) method for the simultaneous
determination of pioglitazone and its metabolites (M-I to M-V) in human serum

urine was developed. The method for serum involved solid-phase and liquid-liquid extraction. Urine with and without enzymic hydrolysis using β -glucuronidase was treated with liquid-liquid extraction. The compds. in the extract were analyzed using HPLC with UV detection at 269 nm. The detection limits of pioglitazone, M-I, M-II, M-III, M-IV and M-V in serum were 0.01-0.05 $\mu g/mL$, those in urine were 0.1-0.5 $\mu g/mL$, and those in urine after enzymic hydrolysis were 0.3-0.5 $\mu g/mL$, resp. The method

was applied to clin. trials of pioglitazone.

IT 111025-46-8, Pioglitazone

RL: ANT (Analyte); ANST (Analytical study)

(high-performance liquid chromatog. determination of pioglitazone and metabolites

in human serum and urine)

RN 111025-46-8 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-(CA INDEX NAME)

PAGE 1-A

PAGE 2-A

Et

IT 101931-00-4 176109-95-8 176109-96-9

176109-97-0

RL: ANT (Analyte); BSU (Biological study, unclassified); MFM (Metabolic formation); ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative)

(high-performance liquid chromatog. determination of pioglitazone and metabolites

in human serum and urine)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

HO₂C-CH₂

L8 ANSWER 22 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1994:524543 CAPLUS

DOCUMENT NUMBER: 121:124543

TITLE: Disposition and metabolism of the hypoglycemic agent

pioglitazone in rats

AUTHOR(S): Krieter, Philip A.; Colletti, Adria E.; Doss, George,

A.; Miller, Randall R.

CORPORATE SOURCE: Department of Drug Metabolism, Merck Research

Laboratories, Rahway, NJ, USA

SOURCE: Drug Metabolism and Disposition (1994), 22(4), 625-30

CODEN: DMDSAI; ISSN: 0090-9556

DOCUMENT TYPE: Journal LANGUAGE: English

AB The disposition and metabolism of [3H]pioglitazone was determined in male rats after oral administration. The peak plasma concentration of 10 $\mu g/mL$ occurred

1 h after dosing at 10 mg/kg p.o.; the apparent plasma terminal half-life was 7.5 h. Most of the radioactivity in plasma ≤8 h after dosing was due to the parent drug. Pioglitazone was highly protein-bound in plasma; only 1-2% was free at concns. of 0.1-10 μg/mL. Within 3 days after oral administration to bile duct-cannulated rats, 36% and 15% of the oral dose was recovered in the bile and urine, resp. The pattern of biliary and urinary metabolites was similar. A total of eight metabolites were isolated and identified on the basis of NMR spectroscopy and MS. Metabolites resulting from hydroxylation of either carbon adjacent to the pyridine ring were conjugated with glucuronic acid or sulfuric acid. The metabolite hydroxylated on the terminal carbon of the Et side chain was

10/520,166

further oxidized to the carboxylic acid derivative Oxidative loss of the terminal carbon led to a nicotinic acid derivative and loss of both carbon atoms to the corresponding 3-hydroxypyridine derivative that was excreted as the sulfate conjugate. The two carboxylic acid metabolites were also conjugated with taurine.

IT 146062-44-4 146062-48-8 157142-90-0 157142-91-1 157142-92-2 157142-93-3 157142-94-4 157142-95-5 186751-40-6 RL: FORM (Formation, nonpreparative)

(formation of, as pioglitazone metabolite)

RN 146062-44-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-[5-(1-hydroxyethyl)-2-pyridinyl]ethoxy]phenyl]methyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

RN 146062-48-8 CAPLUS
CN 3-Pyridineacetic acid, 6-[2-[4-[(2,4-dioxo-5-thiazolidinyl)methyl]phenoxy]
ethyl]- (9CI) (CA INDEX NAME)

| Et

L8 ANSWER 23 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1993:472595 CAPLUS

DOCUMENT NUMBER:

119:72595

TITLE:

Preparation of [[(pyridyl)ethoxy]benzyl]thiazolidinedi

ones as antidiabetics

INVENTOR(S):

Sohda, Takashi; Ikeda, Hitoshi; Greenfield, John C.;

Colca, Jerry R.; Petzold, Edgar N.

PATENT ASSIGNEE(S):

Upjohn Co., USA; Takeda Chemical Industries, Ltd.

SOURCE:

PCT Int. Appl., 43 pp.

DOCUMENT TYPE:

CODEN: PIXXD2 Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9218501	A1 19921029	WO 1992-US2566	19920406
W: AU, BB, BG,	BR, CA, CS, FI,	HU, KP, KR, LK, MG, MN,	MW, NO, PL,
. RO, RU, SD,			
RW: AT, BE, BF,	BJ, CF, CG, CH,	CI, CM, DE, DK, ES, FR,	GA, GB, GN,
GR, IT, LU,	MC, ML, MR, NL,	SE, SN, TD, TG	
CA 2106967	A1 19921012	CA 1992-2106967	19920406
CA 2106967	C 20031209		
AU 9217432	A 19921117	AU 1992-17432	19920406
EP 579733	A1 19940126	EP 1992-910028	19920406

EP 579733	B1	20010620			
R: AT, BE, CH,	DE, DK	, ES, FR,	GB, GR, IT, LI, LU,	MC, N	IL, SE
HU 68370	A2	19950628	HU 1993-2859		19920406
AT 202352	T	20010715	AT 1992-910028		19920406
ES 2161692	T3	20011216	ES 1992-910028		19920406
JP 05086057	A	19930406	JP 1992-85405		19920407
JP 3176694	B2	20010618			
NO 9303615	Α	19931008	NO 1993-3615		19931008
US 5441971	A	19950815	US 1993-137135		19931012
GR 3036623	Т3	20011231	GR 2001-401481		20010917
PRIORITY APPLN. INFO.:			JP 1991-78836	Α	19910411
			WO 1992-US2566	Α	19920406
OMITED COIDCE (C).	MADDAT	110.7250	:		

OTHER SOURCE(S):

MARPAT 119:72595

GI

AB Title compds. I [X = CH2, CO; Q = Ac, MeCHOR, CH2CO2H; R = H, acyl; or Q = Et when X = CO] were prepared as antidiabetics and hypolipemics. Thus, cyclocondensation of Me 2-bromo-3-[4-[2-[5-(1-methoxymethoxyethyl)-2-pyridyl]ethoxy]phenyl]propionate (preparation given) with thiourea followed by deprotection then acetylation, gave title compound I [X = CH2; Q = Ac] (II). II at 0.005 weight% in chow diet for mice gave 56% reduction in blood sugar level

and 43% reduction in plasma lipid level. Tablets containing I were prepared

IT 101931-00-4P 146062-44-4P 146062-45-5P

146062-46-6P 146062-47-7P 146062-48-8P

146062-49-9P 146062-50-2P 146062-51-3P

146062-52-4P 146062-53-5P 146062-54-6P

146062-55-7P 146062-56-8P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of, as antidiabetic)

RN 101931-00-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)-2-

hydroxyethoxy]phenyl]methyl]- (CA INDEX NAME)

| Et

RN 146062-44-4 CAPLUS

CN

2,4-Thiazolidinedione, 5-[[4-[2-[5-(1-hydroxyethy1)-2-pyridiny1]ethoxy]phenyl]methyl]- (CA INDEX NAME)

L8 ANSWER 24 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1991:408602 CAPLUS

DOCUMENT NUMBER:

115:8602

TITLE:

Studies on antidiabetic agents: synthesis and

hypoglycemic activity of 5-[4-(pyridylalkoxy)benzyl]-

2,4-thiazolidinediones

AUTHOR (S):

Kees, Kenneth L.

CORPORATE SOURCE:

Wyeth-Ayerst Res., USA

SOURCE:

Chemtracts: Organic Chemistry (1991), 4(1), 82-6

CODEN: CMOCEI; ISSN: 0895-4445

DOCUMENT TYPE:

Journal; General Review

LANGUAGE:

English

AB The title research of T. Sohda et al. (1990) is reviewed with commentary and 11 refs.

IT 101931-05-9DP, derivs.

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and hypoglycemic activity of)

RN 101931-05-9 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-hydroxy-2-(2-pyridinyl)ethoxy]phenyl]methy 1]- (9CI) (CA INDEX NAME)

ANSWER 25 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1990:423750 CAPLUS

DOCUMENT NUMBER:

TITLE: Studies on antidiabetic agents. Synthesis and

hypoglycemic activity of 5-[4-(pyridylalkoxy)benzyl]-

2,4-thiazolidinediones

AUTHOR (S): Sohda, T.; Momose, Y.; Meguro, K.; Kawamatsu, Y.;

Sugiyama, Y.; Ikeda, H.

CORPORATE SOURCE: Cent. Res. Div., Takeda Chem. Ind. Ltd., Osaka, 532,

Japan

SOURCE: Arzneimittel-Forschung (1990), 40(1), 37-42

CODEN: ARZNAD; ISSN: 0004-4172

DOCUMENT TYPE:

Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 113:23750

GT

The synthesis of a series of title compds. I (R = H, Me, OH; R1 = H, 3-, AB 5-, 6-Me, 5-Et, 6-CH2OH; R2 = H, OH, CH2OH) is described. I were evaluated for hypoglycemic and hypolipidemic activities in genetically obese and diabetic mice. 2-(2-Pyridyl)alkoxy derivs. were found to have much better hypoglycemic and hypolipidemic activities than 2-(3-pyridyl)and 2-(4-pyridyl)alkoxy derivs. or even the previously reported compound, ciglitazone. The introduction of a hydroxyl group at the 2-position of the ethoxy chain potentiated the activities. Among the potent compds.,

Ι

RN 74773-19-6 CAPLUS
CN 2,4-Thiazolidinedione, 5-[[4-[2-(6-methyl-2-pyridinyl)ethoxy]phenyl]methyl
]- (9CI) (CA INDEX NAME)

IT 101930-98-7P 101930-99-8P 101931-00-4P 101931-04-8P 101931-05-9P 101931-06-0P

IT 112529-15-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 112529-15-4 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-(5-ethyl-2-pyridinyl)ethoxy]phenyl]methyl]-

, hydrochloride (1:1) (CA INDEX NAME)

● HCl

L8 ANSWER 26 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1986:186401 CAPLUS

DOCUMENT NUMBER:

104:186401

TITLE:

Thiazolidinedione derivatives, and their medicinal

compositions

INVENTOR(S):

Meguro, Kanji; Fujita, Takeshi

PATENT ASSIGNEE(S):

Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 16 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 8504171	A1	19850926	WO 1984-JP445	19840914

W: MC, US			
JP 60208980	A 19851021 JP 1985-41584		19850301
JP 05070633	B 19931005		
US 4582839	A 19860415 US 1985-711536		19850307
EP 155845	A1 19850925 EP 1985-301895		19850319
R: AT, BE, CH,	DE, FR, GB, IT, LI, LU, NL, SE		
CA 1263961	A1 19891219 CA 1985-476976		19850320
PRIORITY APPLN. INFO.:	WO 1984-US117	A2	19840321
	WO 1984-JP117	Α	19840321
	US 1984-624689	A2	19840611
	WO 1984-JP445	Α	19840914
OMITTE COIMAN (A)	CD CD CD 104 106401 . MAD DATE 104 106401		

OTHER SOURCE(S):

CASREACT 104:186401; MARPAT 104:186401

GI

The title compds. I (R1, R2 = H, alkyl; R3 = H, acyl; n = 0, 1) were prepared Thus, a mixture of 3.2 g the propionate II, 558 mg thiourea, 599 mL NaOAc, and 30 mL EtOH was refluxed for 4 h, 30 mL 6N HCl added, and the resulting mixture refluxed for 16 more hours to give 0.95 g I (R1 = 6-Me, R2 = R3 = H, n = 0). I decreased blood sugar and lipids by 22-53% or 11-58%, resp., in mice. I can be administered in the form of capsules, tablets, powders, etc.

IT 101930-98-7P 101930-99-8P 101931-00-4P 101931-01-5P 101931-02-6P 101931-03-7P 101931-04-8P 101931-05-9P 101931-06-0P 101931-07-1P 101931-08-2P 101946-34-3P

101931-07-1P 101931-08-2P 101946-34-3P

101946-35-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as hypolipemic and hypoglycemic agent)

RN 101930-98-7 CAPLUS

CN 2,4-Thiazolidinedione, 5-[[4-[2-hydroxy-2-(6-methyl-2-pyridinyl)ethoxy]phenyl]methyl]- (9CI) (CA INDEX NAME)